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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO
09/709,785	11/03/2000	Anne N. Murphy	660088.433C1	4280
759	90 10/21/2002			
Stephen J Rosenman PhD 701 Fifth Avenue Suite 6300			EXAMINER	
			CHAKRABARTI, ARUN K	
Seattle, WA 98104-7092			ART UNIT	PAPER NUMBER
			1634	

Please find below and/or attached an Office communication concerning this application or proceeding.

Office Action Summary

Period for Reply

Application No. Applicant(s) 09/709,785

Murphy

Examiner Arun Chakrabarti Art Unit 1634

- The MAILING DATE of this communication appears on the cover sheet with the correspondence address -

	d Trademark Office (Rev. 04-01)	Office Action Summary	Part of Paper No. 17
L		or	
	otice of Dreftsperson's Patent Drawing Review (PTO-948) formetion Disclosure Statement(s) (PTO-1449) Paper No(s),	 Natice of Informal Patent Application (Other; 	PTO-152)
	otice of References Cited (PTO-892)	4) Interview Summary (PTO-413) Paper I	
Attachm			
15)	Acknowledgement is made of a claim for	or domestic priority under 35 U.S.C. §§ 120) and/or 121.
a) [The translation of the foreign language	e provisional application has been received.	
		or domestic priority under 35 U.S.C. § 119('e).
*5		ational Bureau (PCT Rule 17.2(a)). ' a list of the certified copies not received.	
	3. Copies of the certified copies of the	e priority documents have been received in	this National Stage
	2. Certified copies of the priority docu	uments have been received in Application N	lo
}	1. Certified copies of the priority docu	uments have been received.	
a) [☐ All b)☐ Some* c/☐ None of:		
13)	Acknowledgement is made of a claim for	or foreign priority under 35 U.S.C. § 119(a)	-(d) or (f).
1	under 35 U.S.C. §§ 119 and 120		
121	The oath or declaration is objected to by		
	If approved, corrected drawings are requi		
11)		nis: aJ□ approved	
		tion to the drawing(s) be held in abeyance. See	
101	The drawing(s) filed on	is/are a) □ accepted or b) □ objecte	d to by the Examiner.
1	The specification is objected to by the E	Examiner.	
1	ation Papers		
1		are subject to restric	
6) 🔀	Claim(s) 92-107		is/are rejected.
5) 🗆	Claim(s)		is/are allowed.
	4a) Of the above, claim(s) 109-139	is/ar	e withdrawn from consideration.
4) X	Claim(s) 92-139	is/are	pending in the application.
_	ition of Claims		
	closed in accordance with the practice to	allowance except for formal matters, prose under Ex parte Quayle, 1935 C.D. 11; 453	
2a) 💢	This action is FINAL. 2b)	This action is non-final.	
Status	Responsive to communication(s) filed or	n Sep 16, 2002	,
- Any r	reply received by the Office later than three months efter that of patent term adjustment. See 37 CFR 1.704(b).	statute, ceuse the epplication to become ABANDONED (35 U.S meilling date of this communication, even if timely filed, may re-	
- If the	ng date of this communication. Period for reply specified above is less than thirty (30) days, Period for reply is specified above, the meximum statutory p	a reply within the statutory minimum of thirty (30) days will be seriod will apply and will expire SIX (6) MONTHS from the mailin	e considered timely. ng date of this communication.
- Exter	sions of time may be evailable under the provisions of 37 CF	R 1.136 (e). In no event, however, mey a reply be timely filed	after SIX (6) MONTHS from the
	HORTENED STATUTORY PERIOD FOR REA MAILING DATE OF THIS COMMUNICATI	PLY IS SET TO EXPIRE3MONTH	I(S) FROM

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DETAILED ACTION

Specification

 Applicant has amended claims 96,104, 107, and 108 along with several parts of the specification in Paper NO: 15.

Claim Rejections - 35 USC 8 112

- 2. The following is a quotation of the second paragraph of 35 U.S.C. 112:
 - The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.
- Claims 107 is rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

Claim 107 is rejected over the use of the trademarks XPRESS and FLAG. Although the use of trademarks is permissible in patent applications, the proprietary nature of the marks should be respected and every effort made to prevent their use in any manner which might adversely affect their validity as trademarks. Moreover, MPEP 7.35.01 recites, "When a trademark or trade name is used in a claim as a limitation to identify or describe a particular material or product, the claim does not comply with the requirements of 35 U.S.C. 112, second paragraph. See Ex parte Simpson, 218 USPQ 1020 (Bd. App. 1982). The claim scope is uncertain since the trademark or

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trade name cannot be used properly to identify any particular material or product. A trademark or trade name is used to identify a source of goods, and not the goods themselves. Thus, a trademark or trade name does not identify or describe the goods associated with the trademark or trade name." In the present case, the trademark or trade name is used to identify/describe the tags or labels of polypeptides and accordingly, the identification/description is indefinite.

Claim Rejections - 35 USC § 103

- The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all
 obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CAR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103© and potential 35 U.S.C. 102(f) or (g) prior art under 35 U.S.C. 103(a).

Claims 92-94, 97-103 and 105 are rejected under 35 U.S.C. 103 (a) over Marban et al.
 (U.S. Patent 6,183,948 B1) (February 6, 2001) in view of Luban et al. (U.S. Patent 5,773,225)
 (June 30, 1998) further in view of Anderson et al. (U.S. Patent 6,140,067) (October 31, 2000).

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Marban et al teach a method for screening for an agent that alters mitochondrial permeability transition (Abstract), comprising the steps of:

- a) contacting a host cell comprising a mitochondrion with a candidate agent and an inducer of MPT (Abstract and Column 1, line 54 to column 2, line 20 and Claims 1, 17 and 18);
 - b) exposing the cell to an excitation energy (Column 14, lines 8-57);
- c) detecting a level of energy transfer between the first and second energy transfer molecules (Column 14, lines 8-57 and Claims 1, 17 and 18); and
- d) comparing the level of energy transfer to a first reference level generated in the absence of candidate agent, and therefrom identifying an agent that alters MPT (Figure 8 and Claims 1, 17 and 18).

Marban et al teach a method wherein the candidate agent increases or decreases energy transfer between the first and second energy transfer molecules (Figure 8).

Marban et al teach a method for altering survival of a cell and MPT, comprising contacting a mitochondrion with an identified agent, under conditions and for a time sufficient to alter MPT (Examples 1-4).

Marban et al do not teach a method wherein the host cell comprises (I) a first nucleic acid expression construct, comprising a promoter operably linked to a polynucleotide encoding a mitochondrial permeability transition pore component polypeptide fused to a polynucleotide encoding a first energy transfer molecule or a variant thereof, and (ii) a second nucleic expression construct, comprising a promoter operably linked to a polynucleotide encoding a cyclophilin

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polypeptide fused to a polynucleotide encoding a second energy transfer molecule or a variant thereof, wherein binding of the mitochondrial permeability transition pore component polypeptide to the cyclophilin polypeptide results in detectable energy transfer between the first and second energy transfer molecules.

Luban et al. teach a method wherein the host cell comprises (I) a first nucleic acid expression construct, comprising a promoter operably linked to a polynucleotide encoding a first energy transfer molecule or a variant thereof, and (ii) a second nucleic expression construct, comprising a promoter operably linked to a polynucleotide encoding a cyclophilin polypeptide fused to a polynucleotide encoding a second energy transfer molecule or a variant thereof, wherein binding of the polypeptide to the cyclophilin polypeptide results in detectable energy transfer between the first and second energy transfer molecules (Figures 1-2 and Column 6, line 40 to column 9, line 25).

It would have been *prima facie* obvious to one having ordinary skill in the art at the time the invention was made to combine and substitute the cyclophilin D containing fusion protein of Luban et al. in the method for screening an agent that can alter MPT of Marban et al., since Luban et al. state, "The above assay can also be extended to assays using protein expressed in baculovirus, tissue culture cells or Gag purified from virus." (Column 5, lines 22-24). An ordinary practitioner would have been motivated to combine and substitute the cyclophilin D containing fusion nucleic acid construct of Luban et al. in the method for screening an agent that can alter MPT of Marban et al in order to achieve the express advantages, as noted by Luban et

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al., of an invention that can also be extended to assays using protein expressed in baculovirus, tissue culture cells or Gag purified from virus.

Marban et al in view of Luban et al do not teach nucleotide construct encoding a mitochondrial permeability transition pore component polypeptide fused to a polynucleotide wherein binding of the mitochondrial permeability transition pore component polypeptide to the cyclophilin polypeptide results in detectable energy transfer between the first and second energy transfer molecules.

Anderson et al teach a method for detecting an agent that alters mitochondrial permeability transition (Abstract and Claims 1-3 and 87).

Anderson et al teach cyclophilin D polypeptide and adenine nucleotide translocator polypeptide as mitochondrial membrane component which can naturally interact and bind to each other (Claims 23 and 91 and Column 3, lines 49-62).

Anderson et al teach the identification of an agent by comparing the altered mitochondrial function in presence and absence of the candidate agent (Claims 1-3).

Anderson et al teach a method for altering survival of a cell, comprising contacting a cell with an agent under conditions and for a time sufficient to modulate cell survival and alter MPT (Example 1).

Anderson et al do not teach a method of contacting a cyclophilin D polypeptide fusion protein with other protein.

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Luban et al teach a method of contacting a cyclophilin D polypeptide fusion protein with other protein (Claims 1, 2 and 11).

Anderson et al do not teach a method wherein cyclophilin D polypeptide is immobilized on a solid support.

Luban et al teach a method wherein cyclophilin D polypeptide is immobilized on a solid support (Claim 6 and Column 16, lines 30-56).

Anderson et al do not teach a method wherein the fusion protein comprises a ligand for a receptor.

Luban et al teach a method wherein the fusion protein comprises a ligand for a receptor (Claims 7 and 8).

Anderson et al. teach nucleotide construct encoding a mitochondrial permeability transition pore component polypeptide fused to a polynucleotide wherein binding of the mitochondrial permeability transition pore component polypeptide to the cyclophilin polypeptide results in detectable energy transfer between the first and second energy transfer molecules (Claims 23 and 118 and Column 3, line 49 to Column 5, line 22). Moreover, Anderson et al teach cyclophilin D polypeptide and adenine nucleotide translocator polypeptide as mitochondrial membrane component which can naturally interact and bind to each other (Column 3, lines 49-63).

It would have been *prima facie* obvious to one having ordinary skill in the art at the time the invention was made to combine and substitute the mitochondrial permeability transition pore

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component polypeptide of Anderson et al. in the method for screening an agent of Marban et al in view of Luban et al., since Anderson et al. state, "The present invention relates to improved screening assays for therapeutic agents useful in the treatment of type 2 diabetes mellitus, by comparing the levels of one or more indicators of altered mitochondrial function." (Abstract, lines 1-6). By employing scientific reasoning and in order to study the determinants of mitochondrial import which causes alteration of MPT, an ordinary practitioner would have been motivated to combine and substitute the the mitochondrial permeability transition pore component polypeptide of Anderson et al. in the method for screening an agent of Marban et al in view of Luban et al. in order to achieve the express advantages, as noted by Anderson et al., of an invention that relates to improved screening assays for therapeutic agents useful in the treatment of type 2 diabetes mellitus, by comparing the levels of one or more indicators of altered mitochondrial function such as MPT.

Claims 92-94, and 96-105 are rejected under 35 U.S.C. 103 (a) over Marban et al. (U.S. Patent 6,183,948 B1) (February 6, 2001) in view of Luban et al. (U.S. Patent 5,773,225) (June 30, 1998) further in view of Anderson et al. (U.S. Patent 6,140,067) (October 31, 2000). further in view of Briggs et al. (U.S. Patent 6,211,440 B1) (April 3, 2001).

Marban et al. in view of Luban et al. further in view of Anderson et al. teach the method of claims 92-94, 97-103 and 105 as described above.

Marban et al. in view of Luban et al. further in view of Anderson et al. do not teach the method wherein the energy transfer molecules are selected from green fluorescent protein (GFP).

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Briggs et al. teach the method wherein the energy transfer molecules are selected from green fluorescent protein (GFP). (Column 42, lines 39-46).

It would have been *prima facie* obvious to one having ordinary skill in the art at the time the invention was made to combine and substitute the green fluorescent protein of Briggs et al. in the method to identify compounds affecting mitochondrial permeability of Marban et al. in view of Luban et al. further in view of Anderson et al. since Briggs et al. state, "Useful labels in the present invention include biotin for staining with labeled streptavidin conjugate, magnetic beads, fluorescent dyes (e.g., fluorescein, Texas Red, rhodamine, green fluorescence protein, and the like) (Column 42, lines 39-42)". By using the scientific reasoning and strong motivation provided by Briggs et al., an ordinary practitioner would have been motivated to combine and substitute the green fluorescent protein of Briggs et al. in the method to identify compounds affecting mitochondrial permeability of Marban et al in view of Luban et al. further in view of Anderson et al. in order to achieve the express advantages, as noted by Briggs et al., of useful label green fluorescence protein.

7. Claims 92-95, 97-103, and 105-106 are rejected under 35 U.S.C. 103 (a) over Marban et al. (U.S. Patent 6,183,948 B1) (February 6, 2001) in view of Luban et al. (U.S. Patent 5,773,225) (June 30, 1998) further in view of Anderson et al. (U.S. Patent 6,140,067) (October 31, 2000) further in view of Halestrap et al. (Biochimica et Biophysica Acta, (1998), Vol. 1366, pages 79-94).

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Marban et al. in view of Luban et al. further in view of Anderson et al teach the method of claims 92-94, 97-103 and 105 as described above.

Marban et al. in view of Luban et al. further in view of Anderson et al do not teach the method, wherein the human evelophilin A polypeptide is used.

Halestrap et al. teach the method, wherein the human cyclophilin A polypeptide is used (Page 80, Column 2, The molecular mechanism of the MPT Section, Subsection 2.1).

Marban et al. in view of Luban et al. further in view of Anderson et al do not teach the method, wherein the detection reagent is an antibody.

Halestrap et al. teach the method, wherein the detection reagent is an antibody.

It would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to combine and substitute the human cyclophilin A polypeptide and antibody detection reagent of Halestrap et al. in the method for screening an agent that can alter MPT of Marban et al. in view of Luban et al. further in view of Anderson et al., since Halestrap et al. state, "We discuss how the MPT may be involved in determining whether cell death occurs by necrosis or apoptosis. (Abstract, last sentence)". Moreover, Anderson et al teach cyclophilin D polypeptide and adenine nucleotide translocator polypeptide as mitochondrial membrane component which can naturally interact and bind to each other. An ordinary practitioner would have been motivated to combine and substitute the human cyclophilin A polypeptide and antibody detection reagent of Halestrap et al. in the method for screening an agent that can alter MPT of Marban et al. in view of Luban et al. further in view of Anderson et al in order to

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achieve the express advantages, as noted by Anderson et al., of an invention that relates to improved screening assays for therapeutic agents useful in the treatment of type 2 diabetes mellitus, by comparing the levels of one or more indicators of altered mitochondrial function.

And also to elucidate how the MPT may be involved in determining whether cell death occurs by necrosis or apoptosis.

Allowable Subject Matter

8. Claim 108 is objected to as being dependent upon a rejected base claim, but would be allowable if rewritten in independent form including all of the limitations of the base claim and any intervening claims.

Response to Amendment

In response to amendment, 112 (second paragraph) rejection against claims 96, 104, and
 are withdrawn. However, same rejection against claim 107 is hereby properly maintained.
 Rejections under 35 U.S.C. 103 (a) are also properly maintained.

Response to Arguments

10. Applicant's arguments filed on July 30, 2002 and September 16, 2002 have been fully considered but they are not persuasive. In response to applicant's arguments against the references individually, one cannot show nonobviousness by attacking references individually

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where the rejections are based on combinations of references. See *In re Keller*, 642 F.2d 413, 208 USPQ 871 (CCPA 1981); *In re Merck & Co.*, 800 F.2d 1091, 231 USPQ 375 (Fed. Cir. 1986).

Applicant also argues that there is no motivation to combine the references. This argument is not persuasive, especially in the presence of strong motivation provided by Anderson et al. since Anderson et al. state, "The present invention relates to improved screening assays for therapeutic agents useful in the treatment of type 2 diabetes mellitus, by comparing the levels of one or more indicators of altered mitochondrial function." (Abstract, lines 1-6). Same logic is applicable to the motivation of other references cited for 103(a) rejections in the last office action.

Conclusion

 THIS ACTION IS MADE FINAL. Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however,

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will the statutory period for reply expire later than SIX MONTHS from the mailing date of this final action.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Arun Chakrabarti, Ph.D., whose telephone number is (703) 306-5818. If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, W. Gary Jones, can be reached on (703) 308-1152. Any inquiry of a general nature or relating to the status of this application should be directed to the Group analyst Chantae Dessau whose telephone number is (703) 605-1237. Papers related to this application may be submitted to Technology Center 1600 by facsimile transmission via the P.T.O. Fax Center located in Crystal Mall 1. The CM1 Fax Center numbers for Technology Center 1600 are either (703) 305-3014 or (703) 308-4242. Please note that the faxing of such papers must conform with the Notice to Comply published in the Official Gazette, 1096 OG 30 (November 15, 1989).

Arun Chakrabarti Patent Examiner Art Unit 1655 October 17, 2002

> Supervisory Patent Examiner Technology Center 1600